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(54) Title: ENZYMES AS CORROSION INHIBITORS BY REMOVAL OF OXYGEN DISSOLVED IN WATER

(57) Abstract: The invention relates to a new process for water deoxygenation, for application in aerosol products. The process involves the use of an enzymatic system based on an oxidase enzyme a substrate for the oxidase enzyme and catalase. These two enzymes consume oxygen by a two step reaction with the substrate and hydrogen peroxide, which is formed in the first reaction.

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ENZYMES AS CORROSION INHIBITORS BY REMOVAL OF OXYGEN DISSOLVED IN WATER

The invention relates to a new process for water deoxygenation, for application in aerosol products. The
process involves the use of an enzymatic system based on
an oxidase enzyme a substrate for the oxidase enzyme and
catalase. These two enzymes consume oxygen by a two step
reaction with the substrate and hydrogen peroxide, which
is formed in the first reaction.

Corrosion reactions take place in the presence of oxygen, oxygen dissolved in water is responsible for can corrosion in aqueous based formulations within aerosol containers. Currently a number of techniques are used to minimise corrosion in aerosol cans, for example, by the use of corrosion inhibitors or by an oxygen displacement process using nitrogen gas. We have found that corrosion is severely retarded if oxygen is substantially removed from the water present in the aerosol can.

Examples of products found in aerosol cans are air care products, household products, fabric care, waxes, polishes, insecticides, ironing aids, fabric refreshers and carpet cleaners.

The aerosol canister is metal, preferably steel or tin coated steel.

The world market trend is to move towards aerosol formulations containing more water. This is due mainly to regulatory issues: the reduction of the volatile organic

content (VOC) level in aerosol products has involved a reduction in the amount of solvent of many products and an increase in the water content.

When aerosol compositions contain less than 50 ppm of water, corrosion of the aerosol can is not generally a serious problem. However, if the water content is more than 50 ppm in the aerosol composition then corrosion is more likely to occur.

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Many corrosion inhibitor systems have been developed for facing these new regulatory requirements. Examples of these products are borates, benzoates, molybdate, special surfactants (such as sodium lauroyl sarcosinate), sodium nitrite and morpholine and silicates. Usually an acceptable control of the corrosion during the product life of the aerosol canister (around 2 years) is built in to the composition. The above corrosion inhibitors tend to interact with the aerosol canister's surface providing protection against corrosion.

There also can be negative effects of corrosive detinning on the performance of the product. The yellow tin corrosion complex may remain especially when sprayed onto white surfaces. White fabrics or carpets can remain coloured by the liquids of aged aerosol products. Other considerations relate to certain stains like coffee, tea and wine that contain cationic metals. These metals can form brown coloured complexes with tin hydroxyl, causing an evident negative effect of the cleaning formulation onto overall cleaning performance.

Therefore, there is a need to identify better ways to prevent corrosion in aerosol canisters.

Corrosion is an electrochemical process. All corrosion reactions are started by the presence of water and oxygen. Oxygen is a direct participant in the corrosion reaction, acting as a cathode-accepting electron.

Dissolved oxygen present in water based formulations within aerosols is one of the most important factors influencing the rate of corrosion for all metals.

Many corrosion inhibitors have been identified in
the prior art, but none really halt dissolution of the
tin layer in tin-plated aerosol cans over the two years
standard can life, they merely slow it down. Even resin
lacquered tin-plated cans generally need an effective
corrosion inhibitor system.

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T.Godfrey, J.Reichelt: Industrial Enzymology, Nature
Press 1983 - Chapter 4.2: G.Richter - Glucose Oxidase, US
5,980,956, EP 0818960 & EP 0835299 describe the use in
the food industry and especially in canned soft drinks
industry of an enzymatic system based on glucose oxidase
and catalase as an antioxidant primarily to prevent
changes in colour and flavour of foods products both
during processing and in storage.

OUS 4,414,334 describes the use of alcohol oxidase and catalase to remove oxygen dissolved in aqueous

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liquids and discloses the use of such systems in foodstuffs and water distribution systems.

Currently a vacuum process is used to remove oxygen during aerosol product manufacture, which does reduce the oxygen content in the aerosol can. The reduction is only in the aerosol can head space and has little effect on the deoxygenation of the liquid phase. For liquid phase deoxygenation currently used is a method called nitrogen stripping, a process that is quite expensive. The process of the present invention can reduce the oxygen content in the aerosol can during manufacturing and even, during product storage.

We have found that the use of an oxidase enzyme and a substrate for the oxidase enzyme combined with catalase effectively reduces the rate of corrosion in aerosol cans by reducing almost to zero the concentration of oxygen dissolved in the water.

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The process of the invention is particularly
effective at neutral and acidic pH. The deoxygenating
process requires a longer time at alkaline pH: this is
not necessarily a problem since the enzymatic system will
continue to work over time if placed in the aerosol
product.

Other advantages of enzymes are that they are very effective even at low concentration, starting from 0.01 ppm of enzyme and 50 ppm of substrate. The enzymes are also compatible with aerosol formulations and have a low impact on the overall formulation cost.

We present as a feature of the invention an aerosol product comprising a sealed metal canister containing an aerosol composition comprising an oxidase enzyme and a substrate for the enzyme. Preferably catalase is also added.

Alternatively we present as a feature of the invention a method of deoxygenating an aerosol produce comprising filling an aerosol canister with an aerosol composition, an oxidase enzyme and a substrate for the oxidase enzyme and, in any order, filling the aerosol canister with propellant and sealing the aerosol canister.

Preferably a catalase is additionally added into the canister.

Alternatively, we present as a feature of the invention use of an oxidase enzyme and a substrate for the oxidase enzyme as a corrosion inhibiting system for aerosol products. Preferably catalase is also used.

suitable oxidase enzymes are those classified under enzyme classification E.C.1.1.3 (Acting on the CH-OH group of donors with oxygen as acceptor) and include one or more of the following. Not all enzymes produce hydrogen peroxide as a product of the reaction. Therefore in a preferred feature of the invention when such enzymes are used the presence of catalase is not required, for example nucleoside oxidase.

Preferred enzymes are selected from one or more of the following; Malate oxidase, Glucose oxidase, Hexose oxidase, Cholesterol oxidase, Aryl-alcohol oxidase, L-gulonolactone oxidase, Galactose oxidase, Pyranose oxidase, L-sorbose oxidase, Pyridoxine 4-oxidase, Alcohol oxidase, Catechol oxidase, (S)-2-hydroxy-acid oxidase, Ecdysone oxidase, Choline oxidase, Secondary-alcohol oxidase, 4-hydroxymandelate oxidase, Long-chain-alcohol oxidase, Glycerol-3-phosphate oxidase, Xanthine oxidase, Thiamine oxidase, L-galactonolactone oxidase, Cellobiose oxidase, Hydroxyphytanate oxidas, Nucleoside oxidase, N-acylhexosamine oxidase, Polyvinyl-alcohol oxidase, Methanol oxidase, D-arabinono-1,4-lactone oxidase, Vanillyl-alcohol oxidase, Nucleoside oxidase, D-mannitol oxidase and Xylitol oxidase.

A preferred enzyme is Glucose Oxidase. Glucose
Oxidase is a highly specific enzyme derived from the
fungi Aspergillus Niger and Penicillinum. Glucose oxidase
20 is an oxidoreductase, that catalyses the oxidation of DGlucose to gluconic acid using molecular oxygen and
releasing hydrogen peroxide. Glucose oxidase has a
molecular weight of 192000, an optimium temperature of
30-50°C and optimum pH of 4.5-6.5. It is inhibited by
25 heavy metal salts, preferably a chelating agent may be
added to the aerosol composition, and sulfhydyl chelating
agents. The effective amount enzyme needed is from 0.001
ppm to 500 ppm, more preferably between 0.01 and 50 ppm.

Catalase is a common enzyme present in the cell of plants, animals and aerobic bacteria. It promotes the conversion of hydrogen peroxide to water and molecular

oxygen. This reaction is very specific and very fast:
catalase has one of the highest turnover rates for all
enzymes. Catalase is inhibited by urea, freezing and
sunlight under aerobic conditions. The effective amount
of enzyme needed is from 0.001 ppm to 500 ppm, more
preferably between 0.01 and 50 ppm.

The reaction is:

1.
$$2C_6H_{12}O_6 + 2H_2O + 2O_2 \rightarrow 2C_6H_{12}O_7 + 2H_2O_2$$
10 (glucose oxidase)

2.
$$2H_2O_2 \rightarrow 2H_2O + O_2$$
 (catalase)

15 Total reaction:

$$2C_6H_{12}O_6 + O_2 \rightarrow 2C_6H_{12}O_7$$

By forcing the equilibrium of the reaction by an excess of substrate to the oxidase enzyme, it is possible to end up with a final oxygen content close to zero.

Therefore, the concentration of substrate needed in order to increase the velocity of the first reaction is ideally greater than the Km of the enzyme selected (Km is the Michael's constant and is the affinity of the enzyme for the substrate, i.e. the concentration at which 50% of the enzyme binding sites are occupied). Typical Km's are 10^{-1} M to 10^{-6} M.

An important feature of the invention is a substrate for the oxidase enzyme used, this may already be present in the composition to be packaged in the aerosol canister

or it might be added. A preferred substrate is D-glucose.

The performance of the new corrosion inhibitor

system has been evaluated first by measuring the dissolved oxygen reduction (Oxy-meter) in a typical conditions and then by a quick method for the evaluation of corrosion, the jar method, using as fill formulation using tap water treated or not with the enzymatic system on a standard epoxy coated piece of aerosol can.

OXY-METER EVALUATION

A 5L glass beaker is used in this test.

4L of tap water are added into the beaker and warmed to 40°C.

pH of the solution is measured and adjusted to desired value. Dissolved oxygen (DO mg/L) pH and Temperature (°C) are measured through an Oxy-Meter YSI 556 MPS.

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The time zero DO value is collected, D-Glucose is added to the solution and immediately after the enzymatic system is dosed.

The reaction is then followed constantly reading the DO value until it reaches a plateau value.

The system is open, so no control to oxygen intake from the air is considered.

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JAR METHOD:

50 ml glass jars with screw plugs are used in this test.

A round piece of a can is cut and applied on the internal surface of the jar screw plug. A cross is cut by a blade on the can piece in order to simulate possible defects on the can walls.

A poly tetra fluoroethylene gasket is also applied on the plug in order to guarantee a good sealing system. The jar is filled with the testing formula and it is stored in the inverted position to obtain the contact between the liquid formula and the tin plated can piece applied on the plug.

The storage is carried out at different temperature (20°C, 40°C and 50°C) for several days up to 1 month. The storage situation is monitored after 1 day, 1 week, 2 weeks, 1 months and compared to reference can pieces and liquids. The can piece appearance is recorded. A

recording data table with the corresponding corrosion rating is reported below:

Corrosion Rating JM	Can piece appearance
0	No difference from reference
1	Low darkening along the cut lines
2	Darkening along the cut lines
3	Strong darkening on all the can piece area

4	Darkening on all the can piece area
5	Evident darkening on all the can piece area
6	Rust

EXAMPLES:

The liquid phases are typically prepared by mixing D5 Glucose anhydrous to warm 40°C tap water, adjusting the
pH to the desired value and then adding the enzymatic
system to start the de-oxygenation reaction.

Components	Ref 1	Ref 2	Ref 4	Ref 5	Ref 6	Ref 7
	mqq	mqq	ppm	ppm	ppm	ppm
	60	250	500	1000	1000	500
D-Glucose	(0.006%)	(0.025%)	(0.05%)	(0.05%)	(0.1%)	(0.05%)
OxyGo 1500	0.0125	0.05	0.1	0.2	0.2	0.1
Tap Water	to 100%	to 100%	to 100%	to 100%	to 100%	to 100%
PH	7	7.8	7.1	7.1	9.1	4.9

Table 2	
Component	Description of component
D-Glucose	D(+)-Glucose anhydrous >99.5% from Fluka
OxyGo 1500	Glucose Oxidase Enzyme with Catalase side activity from Genencor
NaOH	Sodium Hydroxide, 10% solution
H2SO4	Sulphuric Acid, 9% solution

EXAMPLE RESULTS:

The enzymatic corrosion inhibitor system was tested for all formulations using an Oxy-meter evaluation and for formulation Ref.4, Ref.5, Ref. 6 and Ref.7 using the Jar method. Evaluation of possible residual H2O2, due to slow action of catalase, was done for formulation Ref.4, Ref.5, Ref.6 and Ref.7.

Results:

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Product		DO (n	ng/L)at 40°C			
Product	(Oxy-meter evaluation)					
	Time 0	30'	60′	90′		
Ref 1	4.40	4.03	3.79	3.69		
Ref 2	4.24	4.12	3.42	3.18		
Ref 4	4.31	2.06	1.40	1.10		
Ref 5	4.51	1.05	0.84	0.60		
Ref 6	4.50	2.76	2.01	1.49	·	
Ref 7	3.80	1.55	1.48	1.25	 -	

Product	Corrosion rating							
20°C	1 day	(Jar method) 1 day						:h
	uncut	cut	uncut	cut	uncut	cut	uncut	cut
Ref 4	0	0	0	0	0	0	0	1
Ref 5	0	0	0	0	0	0	0	1
Ref 6	0	0	0	0	0	1	0	1
Ref 7	0	0	0	0	/	1	/	/
Tap Water	0	6	0	6	0	6	1	6

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40°C	1 day		1 week	.,	2 week	S	1 mont	h
	uncut	cut	uncut	cut	uncut	cut	uncut	cut
Ref 4	0	0	0	0	0	1	0	1
Ref 5	0	0	0	0	0	1	0	1
Ref 6	0	0	0	0	0	1	0	1
Ref 7	0	0	0	0	1	1	/	/
Tap Water	0	6	0	6	0	6	2	6

50°C	1 d	1 day		1 week		2 weeks		1 month	
	unc	ut Cut	uncut	cut	uncut	cut	uncut	cut	
Ref 4	0	0	0	0	0	0	0	1	
Ref 5	0	0	0	0	0	1	0	2	
Ref 6	0	0	0	0	0	2	0	2	
Ref 7	0	0	0	0	/	/	/	/	
Tap Wat	er 0	6	0	6	0	6	2	6	

Product		H2O2 %	formation	
	10'	30′	60′	90'
Ref 4	0.01	0.00	0.00	0.00
Ref 5	0.00	0.00	0.00	0.00
Ref 6	0.03	0.00	0.00	0.00
Ref 7	0.01	0.00	0.00	0.00

The above results show that the two enzymatic reactions take place relatively quickly, so the H_2O_2 formed in the first step is consumed in the second step.

CLAIMS

- An aerosol product comprising a sealed metal
 canister containing an aerosol composition
 comprising an oxidase enzyme and a substrate for the enzyme.
- 2. An aerosol product is claimed in claim 1 wherein the aerosol composition additionally comprises catalase.
 - 3. An aerosol product as claimed in either claim 1 or claim 2 wherein the aerosol composition comprises >50 ppm of water.

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- 4. An aerosol product as claimed in either claim 2 or claim 3 wherein the oxidase enzyme is glucose oxidase and the substrate is D-glucose.
- 20 5. A method of deoxygenating an aerosol product comprising filling an aerosol canister with an aerosol composition, an oxidase enzyme and a substrate for the oxidase enzyme and, in any order, filling the aerosol canister with propellant, and sealing the aerosol canister.
 - 6. A method as claimed in claim 5 wherein additionally a catalase is added to into the aerosol canister.
- Just of an oxidase enzyme and a substrate for the oxidase enzyme as a corrosion inhibiting system for aerosol products.
- 8. Use of an oxidase enzyme and a substrate for the oxidase enzyme, as claimed in claim 7, in combination with the catalase.

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According	g to International Patent Classification (IPC) or to both national class	idication and IPC					
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Minimum documentation searched (classification system followed by classification symbols) IPC 7 C09K B65D C12N							
Documen	ntation searched other than minimum documentation to the extent th	al such documents are included in the fields so	earched				
Electronic	c data base consulted during the international search (name of data	base and, where practical, search terms used)				
EPO-I	internal, PAJ, WPI Data						
C. DOCU	IMENTS CONSIDERED TO BE RELEVANT						
Category	° Citation of document, with indication, where appropriate, of the	relevani passages	Relevant to claim No.				
Y	US 3 016 336 A (SCOTT DON ET AL 9 January 1962 (1962-01-09) the whole document	.)	1-8				
Y	US 3 095 307 A (SCOTT DON ET AL 25 June 1963 (1963-06-25) the whole document	1-8					
γ	US 3 686 120 A (JOSEPH W. CREEL 22 August 1972 (1972-08-22) the whole document	1-8					
Y	US 3 723 376 A (CHURCH D,US ET 27 March 1973 (1973-03-27) the whole document	AL)	1-8				
	further documents are listed in the continuation of box C.	Y Patent family members are listed	in annov				
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 Special categories of cited documents: 'A' document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international filing date 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'C' document referring to an oral disclosure, use, exhibition or other means 'T' later document published after the International or priority date and not in conflict with the application or othed and the principle or theory underlying invention 'X' document of particular relevance; the claimed invention involve an inventive step when the document is taken 'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step with one or more other such or other means 			the application but early underlying the claimed invention to be considered to occument is taken alone claimed invention eventive step when the ore other such docu-				
'P' docu	ument published prior to the international filing date but or than the priority date claimed	in the art. '&' document member of the same patent					
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INTERNATIONAL SEARCH REPORT

........ation on patent family members

Interna Application No
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Patent document clied in search report		Publication date		Patent family member(s)	Publication date
US 3016336	A	09-01-1962	NONE		
US 3095307	A	25-06-1963	NONE	ی در بین با در این	
US 3686120	A	22-08-1972	CA GB	964407 A1 1318806 A	18-03-1975 31 - 05-1973
US 3723376	A	27-03-1973	NONE		س پپة شد سر د. ۱۰۰۰ که د. په و ها چه د. س

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